EAST Search History

Ref #	Hits	Search Query	DBs	Default Operator	Plurals	Time Stamp
L1	244	(556/404,560/262,568/11).CCLS.	US-PGPUB; USPAT; USOCR; EPO; JPO; DERWENT	OR	OFF	2007/10/01 08:47
L2	21	l1 and phosphonium	US-PGPUB; USPAT; USOCR; EPO; JPO; DERWENT	OR	OFF	2007/10/01 08:48

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PASSWORD:

TERMINAL (ENTER 1, 2, 3, OR ?):2

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         JUL 02
                 LMEDLINE coverage updated
NEWS
         JUL 02
                SCISEARCH enhanced with complete author names
      3
NEWS
         JUL 02
                CHEMCATS accession numbers revised
NEWS
        JUL 02 CA/CAplus enhanced with utility model patents from China
NEWS
        JUL 16 CAplus enhanced with French and German abstracts
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         JUL 18 CA/CAplus patent coverage enhanced
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        AUG 06 BEILSTEIN updated with new compounds
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NEWS 12
        AUG 06
                FSTA enhanced with new thesaurus edition
NEWS 13 AUG 13
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                 patents
NEWS 14 AUG 20
                 CA/CAplus enhanced with CAS indexing in pre-1907 records
NEWS 15 AUG 27
                 Full-text patent databases enhanced with predefined
                 patent family display formats from INPADOCDB
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                 USPATOLD now available on STN
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                 CAS REGISTRY enhanced with additional experimental
        AUG 28
                 spectral property data
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                 STN AnaVist, Version 2.0, now available with Derwent
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NEWS 21
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                 patents
                 EMBASE, EMBAL, and LEMBASE reloaded with enhancements
NEWS 23
        SEP 24
            19 SEPTEMBER 2007: CURRENT WINDOWS VERSION IS V8.2,
NEWS EXPRESS
              CURRENT MACINTOSH VERSION IS V6.0c(ENG) AND V6.0Jc(JP),
              AND CURRENT DISCOVER FILE IS DATED 19 SEPTEMBER 2007.
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FULL ESTIMATED COST

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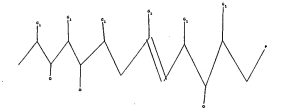
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chain nodes:
1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 19 20 22 23 24 25 chain bonds:
1-2 2-3 2-19 3-4 3-15 4-5 4-20 5-6 5-16 6-7 6-22 7-8 8-9 8-23 9-10 10-11 10-24 11-12 11-17 12-13 12-25 13-14 exact/norm bonds:
2-19 3-15 4-20 5-16 6-22 8-23 10-24 11-17 12-25 exact bonds:
1-2 2-3 3-4 4-5 5-6 6-7 7-8 8-9 9-10 10-11 11-12 12-13 13-14

G1:H,Ak

Match level:

1:CLASS 2:CLASS 3:CLASS 4:CLASS 5:CLASS 6:CLASS 7:CLASS 8:CLASS 9:CLASS 10:CLASS 11:CLASS 12:CLASS 13:CLASS 14:CLASS 15:CLASS 16:CLASS 17:CLASS 19:CLASS 20:CLASS 22:CLASS 23:CLASS 24:CLASS 25:CLASS

L1 STRUCTURE UPLOADED

=> d 11 L1 HAS NO ANSWERS L1 STR

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

Structure attributes must be viewed using STN Express query preparation.

=> s 11

SAMPLE SEARCH INITIATED 08:42:08 FILE 'REGISTRY' SAMPLE SCREEN SEARCH COMPLETED -

BATCH

100.0% PROCESSED

4 ITERATIONS

1 ANSWERS

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS:

ONLINE **COMPLETE**

PROJECTED ITERATIONS:

COMPLETE

4 TO 200

PROJECTED ANSWERS:

1 TO 80

L2

1 SEA SSS SAM L1

=> s l1 full

FULL SEARCH INITIATED 08:42:12 FILE 'REGISTRY'

FULL SCREEN SEARCH COMPLETED - 63 TO ITERATE

100.0% PROCESSED

63 ITERATIONS

14 ANSWERS

SEARCH TIME: 00.00.01

L3

14 SEA SSS FUL L1

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TOTAL

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172.10 172.31

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18 L3

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ANSWER 1 OF 18 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2005:1122416 CAPLUS

DOCUMENT NUMBER:

144:22752

TITLE:

Design, Synthesis, and Biological Evaluation of Potent Discodermolide Fluorescent and Photoaffinity Molecular

Probes

AUTHOR(S):

Smith, Amos B., III; Rucker, Paul V.; Brouard,

Ignacio; Freeze, B. Scott; Xia, Shujun; Horwitz, Susan

Band

CORPORATE SOURCE:

Department of Chemistry, University of Pennsylvania,

Philadelphia, PA, 19104, USA

SOURCE:

Organic Letters (2005), 7(23), 5199-5202

CODEN: ORLEF7; ISSN: 1523-7060

PUBLISHER:

American Chemical Society

DOCUMENT TYPE:

Journal

LANGUAGE:

English

OTHER SOURCE(S):

CASREACT 144:22752

GI

The design, synthesis, and biol. evaluation of a series of (+)-discodermolide mol. probes possessing photoaffinity and fluorescent appendages was achieved. Stereoselective olefin cross-metathesis comprised a key tactic for construction of two of the mol. probes. Three tritium labeled photoaffinity probes I (R = T-4-C6H4-CO-C6H4, R14 = Me, H, R24 = H; R = H, R14 = Me, R24 = T-4-C6H4-CO-4-C6H4CO2CH2) were prepared IT 252342-54-4

Ι

RL: RCT (Reactant); RACT (Reactant or reagent)
 (synthesis, and biol. evaluation of potent discodermolide fluorescent
 and photoaffinity mol. probes)

RN 252342-54-4 CAPLUS

CN Phosphonium, [(2R,3R,4S,5Z,8S,9R,10R,11S,12S,13Z)-3,9-bis[[(1,1-dimethylethyl)dimethylsilyl]oxy]-11-[(4-methoxyphenyl)methoxy]-2,4,6,8,10,12-hexamethyl-5,13,15-hexadecatrienyl]triphenyl-, iodide (9CI) (CA INDEX NAME)

• т-

REFERENCE COUNT:

28 THERE ARE 28 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 2 OF 18 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2005:1080558 CAPLUS

DOCUMENT NUMBER:

144:6608

TITLE:

Design, Synthesis, and Biological Evaluation of

Simplified Analogues of (+)-Discodermolide. Additional

Insights on the Importance of the Diene, the C(7)

Hydroxyl, and the Lactone

AUTHOR(S):

Smith, Amos B., III; Xian, Ming

CORPORATE SOURCE:

Department of Chemistry, Monell Chemical Senses Center, and Laboratory for Research on the Structure of Matter, University of Pennsylvania, Philadelphia,

PA, 19104, USA

SOURCE:

Organic Letters (2005), 7(23), 5229-5232

CODEN: ORLEF7; ISSN: 1523-7060

PUBLISHER:

American Chemical Society

DOCUMENT TYPE:

Journal

LANGUAGE:

English

OTHER SOURCE(S):

CASREACT 144:6608

GΙ

AB The design, synthesis, and biol. evaluation of seven totally synthetic analogs of the antitumor agent (+)-discodermolide are reported. For example, discodermolide analog I (R = H) reacted with methoxymethyl chloride to give I (R = CH2OMe) in 40% yield. Saturation of the terminal diene system, alteration of the substituents on the lactone, and alkylation of the C(7)-hydroxyl group reveal significant structure-activity relationships.

IT 633293-74-0

RL: RCT (Reactant); RACT (Reactant or reagent)
 (preparation of (+)-discodermolide analogs, their antitumor activity, and
 structure-activity relationships)

RN 633293-74-0 CAPLUS

CN Phosphonium, [(2R, 3R, 4S, 5Z, 8S, 9R, 10R, 11S, 12S, 13Z)-9-[[(1,1-dimethylethyl)dimethylsilyl]oxy]-3-(methoxymethoxy)-11-[(4-methoxyphenyl)methoxy]-2,4,6,8,10,12-hexamethyl-5,13,15-hexadecatrienyl]triphenyl-, iodide (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+). Double bond geometry as shown.

• I-

IT 870074-99-0P 870075-28-8P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of (+)-discodermolide analogs, their antitumor activity, and structure-activity relationships)

RN 870074-99-0 CAPLUS

CN Phosphonium, [(2R,3R,4S,5Z,8S,9R,10R,11S,12S)-9-[[(1,1-dimethylethyl)dimethylsilyl]oxy]-3-(methoxymethoxy)-11-[(4-methoxyphenyl)methoxy]-2,4,6,8,10,12-hexamethyl-5-hexadecenyl]triphenyl-, iodide (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-). Double bond geometry as shown.

• I-

RN 870075-28-8 CAPLUS

CN Phosphonium, [(2R, 3R, 4S, 5Z, 8S, 9R, 10R, 11S, 12S, 13Z)-9-[[(1,1-dimethylethyl)dimethylsilyl]oxy]-3-(methoxymethoxy)-11-[(4-methoxyphenyl)methoxy]-2,4,6,8,10,12-hexamethyl-5,13-hexadecadienyl]triphenyl-, iodide (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+). Double bond geometry as shown.

• I-

REFERENCE COUNT:

THERE ARE 28 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 3 OF 18 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2005:312870 CAPLUS

DOCUMENT NUMBER:

143:26411

TITLE:

Toward Understanding How the Lactone Moiety of

Discodermolide Affects Activity

AUTHOR(S):

Shaw, Simon J.; Sundermann, Kurt F.; Burlingame, Mark

A.; Myles, David C.; Freeze, B. Scott; Xian, Ming;

Brouard, Ignacio; Smith, Amos B., III

CORPORATE SOURCE:

Kosan Biosciences, Inc., Hayward, CA, 94545, USA

SOURCE:

Journal of the American Chemical Society (2005),

127(18), 6532-6533

CODEN: JACSAT; ISSN: 0002-7863

American Chemical Society

PUBLISHER:
DOCUMENT TYPE:
LANGUAGE:

Journal English

OTHER SOURCE(S):

CASREACT 143:26411

GΙ

AB A series of simplified discodermolide analogs have been designed and synthesized in an attempt to understand the role of the lactone ring. These synthetic efforts have led to an unsubstituted butyrolactone I being generated, which shows improved activity over the natural product.

IT 633293-74-0

RL: RCT (Reactant); RACT (Reactant or reagent)
(preparation and anticancer activity of discodermolide derivs.)

I

RN 633293-74-0 CAPLUS

CN Phosphonium, [(2R,3R,4S,5Z,8S,9R,10R,11S,12S,13Z)-9-[[(1,1-dimethylethyl)dimethylsilyl]oxy]-3-(methoxymethoxy)-11-[(4-methoxyphenyl)methoxy]-2,4,6,8,10,12-hexamethyl-5,13,15-hexadecatrienyl]triphenyl-, iodide (9CI) (CA INDEX NAME)

I-

REFERENCE COUNT:

THERE ARE 39 CITED REFERENCES AVAILABLE FOR THIS 39 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4ANSWER 4 OF 18 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2004:1141985 CAPLUS

DOCUMENT NUMBER:

142:197745

TITLE:

Design, Synthesis, and Evaluation of Analogues of

(+)-14-Normethyldiscodermolide

AUTHOR(S):

Smith, Amos B., III; Freeze, B. Scott; LaMarche,

Matthew J.; Hirose, Tomoyasu; Brouard, Ignacio; Xian, Ming; Sundermann, Kurt F.; Shaw, Simon J.; Burlingame,

Mark A.; Horwitz, Susan Band; Myles, David C.

CORPORATE SOURCE:

Department of Chemistry, University of Pennsylvania,

Philadelphia, PA, 19104, USA

SOURCE:

Organic Letters (2005), 7(2), 315-318

CODEN: ORLEF7; ISSN: 1523-7060

PUBLISHER:

American Chemical Society

DOCUMENT TYPE:

Journal

LANGUAGE:

English

OTHER SOURCE(S):

CASREACT 142:197745

Ι

GI

AB The design, syntheses, and biol. evaluation of nine totally synthetic analogs of the microtubule-stabilizing agent (+)-14- normethyldiscodermolide (I) are reported. Simplification at the C(21)-C(24) terminal diene and at the C(1)-C(5) lactone moieties reveals significant structure-activity relationships.

IT 835929-84-5P 837383-17-2P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(design, synthesis, and biol. evaluation of analogs of (+)-14-normethyldiscodermolide)

RN 835929-84-5 CAPLUS

CN Phosphonium, [(2R, 3R, 4S, 5Z, 8S, 9R, 10R, 11S, 12S, 13Z)-3, 9-bis[[(1,1-dimethylethyl)dimethylsilyl]oxy]-11-[(4-methoxyphenyl)methoxy]-2,4,8,10,12-pentamethyl-5,13,15-hexadecatrienyl]triphenyl-, iodide (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+). Double bond geometry as shown.

• I-

RN 837383-17-2 CAPLUS

CN Phosphonium, [(2R, 3R, 4S, 5Z, 8S, 9R, 10R, 11S)-3, 9-bis[[(1,1-dimethylethyl)dimethylsilyl]oxy]-11-[(4-methoxyphenyl)methoxy]-2, 4, 8, 10, 12-pentamethyl-5-tridecenyl]triphenyl-, iodide (9CI) (CA INDEX NAME)

REFERENCE COUNT: 38

THERE ARE 38 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 5 OF 18 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2004:1122299 CAPLUS

DOCUMENT NUMBER: 142:197742

TITLE:

Design, Synthesis, and Evaluation of

Carbamate-Substituted Analogues of (+)-Discodermolide

AUTHOR(S):

Smith, Amos B., III; Freeze, B. Scott; LaMarche, Matthew J.; Hirose, Tomoyasu; Brouard, Ignacio; Rucker, Paul V.; Xian, Ming; Sundermann, Kurt F.; Shaw, Simon J.; Burlingame, Mark A.; Horwitz, Susan

Band; Myles, David C.

CORPORATE SOURCE:

Department of Chemistry, University of Pennsylvania,

Philadelphia, PA, 19104, USA

SOURCE:

Organic Letters (2005), 7(2), 311-314

CODEN: ORLEF7; ISSN: 1523-7060

PUBLISHER:

American Chemical Society

DOCUMENT TYPE:

Journal

LANGUAGE:

English

OTHER SOURCE(S):

CASREACT 142:197742

GΙ

- * STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY AVAILABLE VIA OFFLINE PRINT *
- The design, syntheses, and biol. evaluation of 22 totally synthetic AΒ analogs, e.g. I, of the potent microtubule-stabilizing agent (+)-discodermolide (II) have been achieved. Structure-activity relationships of the C(19)-carbamate were defined, exploiting two synthetically simplified scaffolds, as well as the parent (+)-discodermolide framework.
- IT 252342-54-4

RL: RCT (Reactant); RACT (Reactant or reagent) (design, synthesis, and biol. evaluation of carbamate-substituted analogs of (+)-discodermolide)

252342-54-4 CAPLUS RN

Phosphonium, [(2R, 3R, 4S, 5Z, 8S, 9R, 10R, 11S, 12S, 13Z)-3, 9-bis[[(1,1-CN dimethylethyl)dimethylsilyl]oxy]-11-[(4-methoxyphenyl)methoxy]-2,4,6,8,10,12-hexamethyl-5,13,15-hexadecatrienyl]triphenyl-, iodide (9CI) (CA INDEX NAME)

• I-

IT 835929-84-5P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(design, synthesis, and biol. evaluation of carbamate-substituted analogs of (+)-discodermolide)

RN 835929-84-5 CAPLUS

CN Phosphonium, [(2R,3R,4S,5Z,8S,9R,10R,11S,12S,13Z)-3,9-bis[[(1,1-dimethylethyl)dimethylsilyl]oxy]-11-[(4-methoxyphenyl)methoxy]-2,4,8,10,12-pentamethyl-5,13,15-hexadecatrienyl]triphenyl-, iodide (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+). Double bond geometry as shown.

• I-

REFERENCE COUNT:

38 THERE ARE 38 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 6 OF 18 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2004:1016004 CAPLUS

DOCUMENT NUMBER:

142:6360

TITLE:

Synthetic techniques and intermediates for polyhydroxy

dienyl lactones and mimics thereof

INVENTOR(S):

Myles, David C.; Burlingame, Mark; Shaw, Simon James;

Sundermann, Kurt F.; Freeze, Brian Scott; Martin, Ignacio Brouard; Hirose, Tomoyasu; Smith, Amos B.

PATENT ASSIGNEE(S):

The Trustees of the University of Pennsylvania, USA

SOURCE: PCT Int. Appl., 50 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

LANGUAGE:

Patent English

1

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

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                                              WO 2004-US10272
                                                                       20040402
     WO 2004101508
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             ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE, SI,
             SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN,
             TD, TG
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                                                                       20030606
OTHER SOURCE(S):
                          CASREACT 142:6360; MARPAT 142:6360
GΙ
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* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *
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AB Synthetic methods and intermediates, e.g., I·X- [R0 = C1-6-alkyl, C2-6-alkenyl, C2-6-alkynyl, (CH2)r(C3-6-cycloalkyl), CH2-aryl, CH2-heterocycle; r = 0 - 4; R1, R2, R3, R6, R7, R8 = H, C1-10-alkyl; R4 = acid-labile OH protecting group; R5 = oxidatively-labile OH protecting group; R9 = C6-14-aryl; Q = H, acid-labile OH protecting group; (whereby the acid-labile OH protecting group has a mass of 135 Daltons or less and is unbranched at the atom bonded to O of the protected OH); X = halogen], useful in the preparation of lactone containing compds. such as discodermolide and

compds. which mimic the chemical or biol. activity of discodermolide are provided. The synthetic method comprises reaction of halide II with phosphine P(R9)3 for a time and under conditions sufficient to prepare I·X- (whereby the pressure is less than about 10,000 psi). Thus, I·X- [R0 = CH:CHCH:CH2-(Z), R1 = R2 = R3 = R6 = R7 = R8 = Me, R4 = CH2C6H4OMe-4, R5 = Q = SiMe2CMe3, R9 = Ph, X = I] was prepared and used to synthesize (+)-discodermolide (III) via Wittig reaction with aldehyde IV.

633293-74-0P
RL: IMF (Industrial manufacture); RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
 (preparation and Wittig reaction of, with (oxotetrahydropyranyl)propanal derivative; synthetic techniques and intermediates for discodermolide and other polyhydroxy dienyl lactones and mimics thereof)

RN 633293-74-0 CAPLUS

ΙT

CN Phosphonium, [(2R,3R,4S,5Z,8S,9R,10R,11S,12S,13Z)-9-[[(1,1-dimethylethyl)dimethylsilyl]oxy]-3-(methoxymethoxy)-11-[(4-methoxyphenyl)methoxy]-2,4,6,8,10,12-hexamethyl-5,13,15-hexadecatrienyl]triphenyl-, iodide (9CI) (CA INDEX NAME)

I-

T.4 ANSWER 7 OF 18 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2004:303318 CAPLUS

DOCUMENT NUMBER:

141:54112

TITLE:

Design, synthesis and cytotoxicity of 7-deoxy aryl

discodermolide analogues

AUTHOR(S):

Burlingame, Mark A.; Shaw, Simon J.; Sundermann, Kurt F.; Zhang, Dan; Petryka, Joseph; Mendoza, Esteban; Liu, Fenghua; Myles, David C.; LaMarche, Matthew J.; Hirose, Tomoyasu; Freeze, B. Scott; Smith, Amos B.

CORPORATE SOURCE:

Department of Chemistry, Kosan Biosciences Inc.,

Hayward, CA, 94545, USA

SOURCE:

Bioorganic & Medicinal Chemistry Letters (2004),

14(9), 2335-2338

CODEN: BMCLE8; ISSN: 0960-894X

PUBLISHER:

Elsevier Science B.V.

DOCUMENT TYPE:

Journal

LANGUAGE:

English

OTHER SOURCE(S):

CASREACT 141:54112

A series of 7-deoxy discodermolide analogs in which the lactone fragment C' was replaced by aryl substituents were designed, synthesized, and evaluated for cytotoxicity.

IT 252342-54-4

RL: RCT (Reactant); RACT (Reactant or reagent)

(design, synthesis and antitumor cytotoxicity of 7-deoxy aryl

discodermolide analogs)

RN 252342-54-4 CAPLUS

CN Phosphonium, [(2R, 3R, 4S, 5Z, 8S, 9R, 10R, 11S, 12S, 13Z)-3, 9-bis[[(1,1dimethylethyl)dimethylsilyl]oxy]-11-[(4-methoxyphenyl)methoxy]-2,4,6,8,10,12-hexamethyl-5,13,15-hexadecatrienyl]triphenyl-, iodide (9CI) (CA INDEX NAME)

• I-

REFERENCE COUNT:

THERE ARE 37 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 8 OF 18 CAPLUS COPYRIGHT 2007 ACS on STN

37

ACCESSION NUMBER:

2003:810303 CAPLUS

DOCUMENT NUMBER:

140:27700

TITLE:

A Practical Improvement, Enhancing the Large-Scale Synthesis of (+)-Discodermolide: A Third-Generation

Approach

AUTHOR(S):

Smith, Amos B.; Freeze, B. Scott; Brouard, Ignacio;

Hirose, Tomoyasu

CORPORATE SOURCE:

Department of Chemistry, University of Pennsylvania,

Philadelphia, PA, 19104, USA

SOURCE:

Organic Letters (2003), 5(23), 4405-4408

CODEN: ORLEF7; ISSN: 1523-7060 American Chemical Society

PUBLISHER:

Journal

DOCUMENT TYPE: LANGUAGE:

English

OTHER SOURCE(S):

CASREACT 140:27700

GΙ

AB A significant improvement to the Penn one-gram synthesis of (+)-discodermolide has been achieved. Specifically, reduction of the steric bulk of the C(11) hydroxyl protecting group permits formation of the requisite AB Wittig salt I at the expense of the undesired intramol. cyclization upon treatment with PPh3 at ambient pressure.

IT 633293-74-0P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(large-scale synthesis of (+)-discodermolide, a third-generation approach)

RN 633293-74-0 CAPLUS

CN Phosphonium, [(2R,3R,4S,5Z,8S,9R,10R,11S,12S,13Z)-9-[[(1,1-dimethylethyl)dimethylsilyl]oxy]-3-(methoxymethoxy)-11-[(4-methoxyphenyl)methoxy]-2,4,6,8,10,12-hexamethyl-5,13,15-hexadecatrienyl]triphenyl-, iodide (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+). Double bond geometry as shown.

• I-

REFERENCE COUNT:

THERE ARE 26 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 9 OF 18 CAPLUS COPYRIGHT 2007 ACS on STN

26

ACCESSION NUMBER:

2003:500684 CAPLUS

DOCUMENT NUMBER:

139:381288

TITLE:

Synthesis and biological assessment of simplified analogues of the potent microtubule stabilizer

(+)-Discodermolide

AUTHOR(S):

Minguez, Jose M.; Kim, Sun-Young; Giuliano, Kenneth A.; Balachandran, Raghavan; Madiraju, Charitha; Day,

Billy W.; Curran, Dennis P.

CORPORATE SOURCE:

Department of Chemistry, Chevron Science Center,

Pittsburgh, PA, 15260, USA

SOURCE:

Bioorganic & Medicinal Chemistry (2003), 11(15),

3335-3357

CODEN: BMECEP; ISSN: 0968-0896

PUBLISHER:
DOCUMENT TYPE:

Elsevier Science Ltd.

DOCOMENT TIFE

Journal

LANGUAGE:

English

OTHER SOURCE(S):

CASREACT 139:381288

GI

AB An efficient, convergent and stereocontrolled synthesis of simplified analogs (e.g. I) of the potent antimitotic agent (+)-discodermolide has

been achieved and several small libraries have been prepared In all the libraries, the discodermolide Me groups at C14 and C16 and the C7 hydroxy group were removed and the lactone was replaced by simple esters. Other modifications introduced in each series of analogs were related to C11, C17 and C19 of the natural product. Key elements of the synthetic strategy included (a) elaboration of the main subunits from a common intermediate and (b) fragment couplings using Wittig reactions to install the (Z)-olefins. Library components were analyzed for microtubule-stabilizing actions in vitro, for displacement of [3H]paclitaxel from its binding site on tubulin, for antiproliferative activity against human carcinoma cells, and for cell signaling and mitotic spindle alterations by a multiparameter fluorescence cell-based screening technique. The results show that even significant structural simplification can lead to analogs with actions related to microtubule targeting.

IT 623926-76-1P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(synthesis and biol. activity of discodermolide analogs)

RN 623926-76-1 CAPLUS

CN Phosphonium, [(2R,3S,4S,5Z,9R,10R,11S,12S,13Z)-9-[[(1,1-dimethylethyl)dimethylsilyl]oxy]-3,11-bis[(4-methoxyphenyl)methoxy]-2,4,10,12-tetramethyl-5,13,15-hexadecatrienyl]triphenyl-, iodide (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

• I-

PAGE 1-B

[→] OMe

REFERENCE COUNT:

THERE ARE 48 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 10 OF 18 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2003:133020 CAPLUS

138:170004

DOCUMENT NUMBER:

TITLE:

Preparation of compounds which mimic the chemical and

biological properties of discodermolide

INVENTOR(S):

Smith, Amos B., III; Beauchamp, Thomas J.; Lamarche,

Matthew J.; Rucker, Paul

PATENT ASSIGNEE(S):

SOURCE:

The Trustees of the University of Pennsylvania, USA PCT Int. Appl., 333 pp.

PCT Int. Appl., 333 pp. CODEN: PIXXD2

DOCUMENT TYPE:

LANGUAGE:

Patent English

FAMILY ACC. NUM. COUNT:

Г	:	1

PATENT INFORMATION:

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	WO	2003	0135	02		A1	- ,	2003	0220		uo Wo	2002-	US24	 932		2	- -	 806
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	IN	A 2004000974 N 2004KN00289			Α		2006	0331								0040		
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	PRIORITY APPLN. INFO.:										2002-0					0020		
OMITTO	n ~ .		/ ~ \															

OTHER SOURCE(S):

MARPAT 138:170004

GΙ

AB Discodermolide analogs, such as I [R = H, OR33; X = H2, O; R4, R9, R33 = H, acid labile protecting group; R25 = H, carbamoyl, thiocarbamoyl, oxidatively labile protecting group; R16, R32 = H, alkyl], were prepared Synthetic routes to both (-)- and (+)-discodermolide were presented. prepared discodermolide analogs were assayed in vitro for tubulin polymerization

inhibition and for cytotoxicity against human A459 cancer cells.

IT 252342-54-4P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of compds. which mimic the chemical and biol. properties of discodermolide for pharmaceutical use as anticancer agents)

RN

252342-54-4 CAPLUS
Phosphonium, [(2R,3R,4S,5Z,8S,9R,10R,11S,12S,13Z)-3,9-bis[[(1,1-CNdimethylethyl)dimethylsilyl]oxy]-11-[(4-methoxyphenyl)methoxy]-2,4,6,8,10,12-hexamethyl-5,13,15-hexadecatrienyl]triphenyl-, iodide (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+). Double bond geometry as shown.

REFERENCE COUNT:

3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 11 OF 18 CAPLUS COPYRIGHT 2007 ACS on STN L4ACCESSION NUMBER: 2002:575783 CAPLUS DOCUMENT NUMBER: 137:125048 TITLE: Preparation of compounds which mimic the chemical and biological properties of discodermolide INVENTOR(S): Smith, Amos B., III; Beauchamp, Thomas J.; Lamarche, Matthew J. The Trustees of The University of Pennsylvania, USA PATENT ASSIGNEE(S): U.S. Pat. Appl. Publ., 127 pp., Cont.-in-part of U.S. Ser. No. 455,649. SOURCE: CODEN: USXXCO Patent DOCUMENT TYPE: LANGUAGE: English FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

	PATENT NO. US 2002103387				KIN	D	DATE			APPLICATION NO.					DATE				
		2002 6870		87		A1 B2		2002 2005			US 2	2000-	7309	29		2	0001	206	
		5789				A		1998			US 1	.996-	7598	17		1	9961	203	
	US	6031	133			Α		2000	0229		US 1	998-	2187	8		1	9980	211	
	US	6242	616			В1		2001	0605		US 1	999-	4556	49		1	9991	207	
		2431				A1		2002	0613		CA 2	001-	2431	045		2	0011	206	
	WO	2002	0461	50		A2		2002	0613		WO 2	001-	US47	958		2	0011	206	
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		1585		5		A2		2002				002-					0011		
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		14.		FI,			DI,	ĽО,	ĖΚ,	GD,	GR,	11,	тт,	ъυ,	ИL,	SE,	MC,	PT,	
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				NE,	SN,	•													
DD75		2007				A1		2007	0222			006-					0060.		
PRIO	ктлу	APP:	LN.	INFO	. :							996-				A2 19			
									•	US 1998-21878					A2 19				
										US 1999-455649					A2 19991207				

US 1998-121551

A2 19980723

AU 1999-52190 A3 19990720 US 2000-730929 A 20001206 WO 2001-US47958 W 20011206 US 2004-779049 A 20040213

OTHER SOURCE(S): GI

MARPAT 137:125048

AB Discodermolide analogs, such as I [R = H, OR33; X = H2, O; R4, R9, R33 = H, acid labile protecting group; R25 = H, oxidatively labile protecting group; R16, R32 = H, alkyl], were prepared Synthetic routes to both (-)-and (+)-discodermolide were presented.

Ι

IT 252342-54-4P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of compds. which mimic the chemical and biol. properties of discodermolide)

RN 252342-54-4 CAPLUS

CN Phosphonium, [(2R,3R,4S,5Z,8S,9R,10R,11S,12S,13Z)-3,9-bis[[(1,1-dimethylethyl)dimethylsilyl]oxy]-11-[(4-methoxyphenyl)methoxy]-2,4,6,8,10,12-hexamethyl-5,13,15-hexadecatrienyl]triphenyl-, iodide (9CI) (CA INDEX NAME)

• r-

REFERENCE COUNT:

THERE ARE 50 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 12 OF 18 CAPLUS COPYRIGHT 2007 ACS on STN

50

ACCESSION NUMBER:

2002:449643 CAPLUS

DOCUMENT NUMBER:

137:33164

TITLE:

Preparation of compounds which mimic the chemical and

biological properties of discodermolide

INVENTOR(S):

Smith, Amos B., III; Beauchamp, Thomas J.; Lamarche,

Matthew J.

PATENT ASSIGNEE(S):

The Trustees of the University of Pennsylvania Center

for Technology Transfer, USA

SOURCE:

PCT Int. Appl., 267 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

6

PATENT INFORMATION:

PA:	TENT	NO.			KIN	CIND DATE				APPL	ICAT		DATE				
	2002 2002									WO 2	001-	us47	958		2	0011	206
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							MD,										•
US	2002	1033	87		A1		2002	0801	1	US 2	000-	7309	29		2	0001	206
US	6870	058			В2		2005	0322									
CA	2431	045			A1		2002	0613		CA 2	001-	2431	045		2	0011	206
	2002																
EP	1585	725			A2		2005	1019		EP 2	001-	9962	31		21	0011	206
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			FI,							·	•	•	•	•	•		,
ΑU	2002	3004	72		A1		2003	0213		AU 2	002-	3004	72		21	0020	730
IN	2003	KN00	715 .		Α		2005	1202			003-1					0030	
WO	2005	0793	78		A2		2005	0901	1	WO 2	005-1	JS46	43			0050	
WO	2005	0793	7.8		A 3		2006	0216							_		

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             CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD,
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             NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY,
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             AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK,
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             RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML,
             MR, NE, SN, TD, TG
PRIORITY APPLN. INFO.:
                                            US 2000-730929
                                                                 A 20001206
                                            US 1996-759817
                                                                 A2 19961203
                                            US 1998-21878
                                                                 A2 19980211
                                            AU 1999-52190
                                                                 A3 19990720
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                                                                    20011206
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                                                                 Α
                                                                    20040213
OTHER SOURCE(S):
                         MARPAT 137:33164
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X.

GI

AB Discodermolide analogs, such as I [R = H, OR33; X = H2, O; R4, R9, R33 = H, acid labile protecting group; R25 = H, oxidatively labile protecting group; R16, R32 = H, alkyl], were prepared Synthetic routes to both (-)-and (+)-discodermolide were presented.

IT 252342-54-4P

Ι

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of compds. which mimic the chemical and biol. properties of discodermolide)

RN 252342-54-4 CAPLUS

CN Phosphonium, [(2R,3R,4S,5Z,8S,9R,10R,11S,12S,13Z)-3,9-bis[[(1,1-dimethylethyl)dimethylsilyl]oxy]-11-[(4-methoxyphenyl)methoxy]-2,4,6,8,10,12-hexamethyl-5,13,15-hexadecatrienyl]triphenyl-, iodide (9CI) (CA INDEX NAME)

) I-

136:183657

ANSWER 13 OF 18 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: DOCUMENT NUMBER:

2002:123244 CAPLUS

TITLE:

Process for the biomediated preparation of

intermediates for use in the synthesis of polyketides,

... such as epothilone D and discodermolide

INVENTOR(S):

Santi, Daniel V.; Ashley, Gary; Myles, David C. Kosan Biosciences, Inc., USA

PATENT ASSIGNEE(S):

PCT Int. Appl., 129 pp.

SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE: LANGUAGE:

Patent English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PAT	CENT I	NO.			KIN	D	DATE		į	APPL	ICAT	ION 1	NO.		D	ATE	
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							MD,										
							SI,										
			VN,												•	•	•
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			YU,														
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AU	2001	0832	75 ု		A5		20020	0218	Ï	AU 20	001-8	3327	5		20	00108	309

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                                             US 2000-207331P
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                                                                     20000530
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                                                                 W
                                                                     20010529
                                             WO 2001-US25112
                                                                 W
                                                                     20010809
OTHER SOURCE(S):
                         CASREACT 136:183657; MARPAT 136:183657
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GT

The present invention relates to compds., such as I, made by a subset of AB modules from one or more polyketide synthase ("PKS") genes that are used as starting material in the chemical synthesis of novel mols., particularly naturally occurring polyketides or derivs. thereof. The biol. derived intermediates ("bio-intermediates") generally represent particularly difficult compds. to synthesize using traditional chemical approaches due to one or more stereocenters. In one aspect of the invention, an intermediate in the synthesis of epothilone is provided that feeds into the synthetic protocol of Danishefsky and co-workers. In another aspect of the invention, intermediates in the synthesis of discodermolide are provided that feed into the synthetic protocol of Smith and co-workers. By taking advantage of the inherent stereochem. specificity of biol. processes, the syntheses of key intermediates and thus the overall syntheses of compds. like epothilone and discodermolide are greatly simplified.

IT 252342-54-4P

RL: BMF (Bioindustrial manufacture); BPN (Biosynthetic preparation); IMF (Industrial manufacture); RCT (Reactant); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent) (process for the biomediated preparation of intermediates for use in the synthesis of polyketides, such as epothilone D and discodermolide) 252342-54-4 CAPLUS

RN 252342-54-4 CAPLUS
CN Phosphonium, [(2R,3R,4S,5Z,8S,9R,10R,11S,12S,13Z)-3,9-bis[[(1,1-dimethylethyl)dimethylsilyl]oxy]-11-[(4-methoxyphenyl)methoxy]-2,4,6,8,10,12-hexamethyl-5,13,15-hexadecatrienyl]triphenyl-, iodide (9CI) (CA INDEX NAME)

• I-

135:19496

L4 ANSWER 14 OF 18 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: DOCUMENT NUMBER:

2001:412212 CAPLUS

TITLE:

Preparation of intermediates for the synthesis of

discodermolides and their polyhydroxy dienyl lactone

derivatives for pharmaceutical use

INVENTOR(S):

Smith, Amos B., III; Beauchamp, Thomas J.; Lamarche,

Matthew J.; Arimoto, Hirokazu

PATENT ASSIGNEE(S):

The Trustees of the University of Pennsylvania, USA

SOURCE:

U.S., 126 pp., 6096904 Cont.-in-part of U.S.

6,096,904.

CODEN: USXXAM

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PAT	CENT	NO.			KIN		DATE			APPL	ICAT:	ION :	NO.		D	ATE	
US	6242	616			В1		2001	0605		US 1	999-	4556	49		1	9991	207
US	5789	605			Α		1998	0804		US 1	996-	7598	17		1	9961	203
US	6031	133			Α		2000	0229			998-					9980	211
US	6096	904			Α		2000	0801		US 1	998-	1215	51		1	9980	723
	2393				A1		2001	0614		CA 2	000-	2393	968		2	0001	206
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							GB,									TR,	BF,
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	2002									US 2	000	7309	29		2	0001	206
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	2005						2005									0040	
WO	2005	0793	78		A2		2005	0901	1	WO 2	005-1	US46	43		20	0050	211

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WO 2005079378
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             EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LT, LU, MC, NL, PL, PT,
             RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GO, GW, ML,
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PRIORITY APPLN. INFO.:
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                                                                 Al 19961203
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                                                                 Al 19980211
                                             US 1998-121551
                                                                 A2 19980723
                                             AU 1999-52190
                                                                 A3 19990720
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                                             US 2000-730929
                                                                 A1 20001206
                                             WO 2000-US32996
                                                                    20001206
                                             US 2004-779049
                                                                 Α
                                                                    20040213
OTHER SOURCE(S):
                         CASREACT 135:19496; MARPAT 135:19496
GΙ
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Preparation of intermediates, such as I [R11, R12 = alkyl; R14, R15 = acid labile protecting groups; R16 = H, alkyl] and II [R1, R2, R7, R8 = alkyl; R3, R6, R16 = H, alkyl; R4, R9 = acid labile hydroxyl protecting group; R25 = oxidatively labile hydroxyl protecting group; X = :C(J)R16, a Wittig olefination formed from a pyranylalkyl ketone, such as I and II (X = P+Ph3I-)], for the synthesis of discodermolides and their analogs, which are useful as pharmaceuticals, was presented. Thus, synthon III (R14 = R15 = SiMe2CMe3) was prepared via a multistep synthetic sequence starting from (2R)-3-hydroxy-2-methylpropanoic acid Me ester. The synthetic utility of II was subsequently demonstrated by its use in the preparation of (-)-discodermolide.

IT 252342-54-4P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of intermediates for the synthesis of discodermolides and their polyhydroxy dienyl lactone derivs. for pharmaceutical use)

RN 252342-54-4 CAPLUS

CN

Phosphonium, [(2R,3R,4S,5Z,8S,9R,10R,11S,12S,13Z)-3,9-bis[[(1,1-dimethylethyl)dimethylsilyl]oxy]-11-[(4-methoxyphenyl)methoxy]-2,4,6,8,10,12-hexamethyl-5,13,15-hexadecatrienyl]triphenyl-, iodide (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+). Double bond geometry as shown.

♠ T =

REFERENCE COUNT:

THERE ARE 30 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 15 OF 18 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2000:597937 CAPLUS 133:335118

DOCUMENT NUMBER: TITLE:

Evolution of a Gram-Scale Synthesis of

(+)-Discodermolide

AUTHOR(S):

Smith, Amos B., III; Beauchamp, Thomas J.; LaMarche, Matthew J.; Kaufman, Michael D.; Qiu, Yuping; Arimoto,

Hirokazu; Jones, David R.; Kobayashi, Kaoru

CORPORATE SOURCE:

Department of Chemistry Monell Chemical Senses Center

and Laboratory for Research on the Structure of

Matter, University of Pennsylvania, Philadelphia, PA,

19104, USA

SOURCE:

Journal of the American Chemical Society (2000),

122(36), 8654-8664

CODEN: JACSAT; ISSN: 0002-7863

PUBLISHER:

American Chemical Society

DOCUMENT TYPE:

Journal

LANGUAGE:

English

OTHER SOURCE(S):

CASREACT 133:335118

GI

AB An efficient, highly convergent, stereocontrolled total synthesis of the potent antimitotic agent (+)-discodermolide (I) has been achieved on gram scale. Key elements of the successful strategy include (1) elaboration of three advanced fragments from a common precursor (CP) which embodies the repeating stereochem. triad of the discodermolide backbone, (2) \(\sigma\)-bond installation of the Z trisubstituted olefin, exploiting a modified Negishi cross-coupling reaction, (3) synthesis of a late-stage phosphonium salt utilizing high pressure, and (4) Wittig installation of the Z disubstituted olefin and the terminal (Z)-diene.

Ι

IT 252342-54-4P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(evolution of a gram-scale synthesis of (+)-discodermolide)

RN 252342-54-4 CAPLUS

CN Phosphonium, [(2R, 3R, 4S, 5Z, 8S, 9R, 10R, 11S, 12S, 13Z)-3, 9-bis[[(1,1-dimethylethyl)dimethylsilyl]oxy]-11-[(4-methoxyphenyl)methoxy]-2,4,6,8,10,12-hexamethyl-5,13,15-hexadecatrienyl]triphenyl-, iodide (9CI) (CA INDEX NAME)

L4 ANSWER 16 OF 18 CAPLUS COPYRIGHT 2007 ACS on STN

133:135166

ACCESSION NUMBER:

2000:531688 CAPLUS

DOCUMENT NUMBER: TITLE:

Preparation of intermediates for the synthesis of

discodermolides and their polyhydroxy dienyl lactone

derivatives for pharmaceutical use

INVENTOR(S):

Smith, Amos B., III; Qiu, Yuping; Kaufman, Michael; Arimoto, Hirokazu; Jones, David R.; Kobayashi, Kaoru;

Beauchamp, Thomas J.

PATENT ASSIGNEE(S):

The Trustees of the University of Pennsylvania, USA U.S., 83 pp., Cont.-in-part of U.S. 5,789,605.

SOURCE:

CODEN: USXXAM

DOCUMENT TYPE:

Patent English

LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

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	2002		72		A1		2003				002-					0020		
	2005				A1		2005		,		004-					0040		
	2005				A2		2005				005-1					0050		
	2005				A3		2006		'	" 2	005	0540	43		. 2	0030	Z I I	
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							DE,											
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AB Preparation of intermediates, such as I [R11, R12 = alkyl; R14, R15 = acid labile protecting groups; R16 = H, alkyl], for the synthesis of discodermolides and their analogs, which are useful as pharmaceuticals, was presented. Thus, synthon II (R14 = R15 = SiMe2CMe3) was prepared via a multistep synthetic sequence starting from (2R)-3-hydroxy-2-methylpropanoic acid Me ester. The synthetic utility of II was subsequently demonstrated by its use in the preparation of (-)-discodermolide. IT 252342-54-4P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of intermediates for the synthesis of discodermolides and their polyhydroxy dienyl lactone derivs. for pharmaceutical use)

RN 252342-54-4 CAPLUS

CN Phosphonium, [(2R,3R,4S,5Z,8S,9R,10R,11S,12S,13Z)-3,9-bis[[(1,1-dimethylethyl)dimethylsilyl]oxy]-11-[(4-methoxyphenyl)methoxy]-2,4,6,8,10,12-hexamethyl-5,13,15-hexadecatrienyl]triphenyl-, iodide (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+). Double bond geometry as shown.

Ι

● T-

REFERENCE COUNT:

31 THERE ARE 31 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 17 OF 18 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2000:84572 CAPLUS

DOCUMENT NUMBER:

132:137207

TITLE:

Preparation of intermediates for the synthesis of discodermolides and their polyhydroxy dienyl lactone

derivatives for pharmaceutical use

Smith, Amos B., III; Qiu, Yuping; Kaufman, Michael; Arimoto, Hirokazu; Jones, David R.; Kobayashi, Kaoru;

Beauchamp, Thomas J.

PATENT ASSIGNEE(S):

SOURCE:

The Trustees of the University of Pennsylvania, USA PCT Int. Appl., 201 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

INVENTOR(S):

LANGUAGE:

Patent English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PAT	CENT	NO.			KIN:	D	DATE			APPL	ICAT	ION	NO.		D.	ATE		
	2000 2000 W:	0048			A2 A3			0203 0921		WO 1	999-	 US16	 369		1	9990	720	
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OTHER SOURCE(S):

GI

II

MARPAT 132:137207

Preparation of intermediates, such as I [R11, R12 = alkyl; R14, R15 = acid AB labile protecting groups; R16 = H, alkyl], for the synthesis of discodermolides and their analogs, which are useful as pharmaceuticals, was presented. Thus, synthon II (R14 = R15 = SiMe2CMe3) was prepared via a multistep synthetic sequence starting from (2R)-3-hydroxy-2methylpropanoic acid Me ester. The synthetic utility of II was subsequently demonstrated by its use in the preparation of (-)-discodermolide. IT 252342-54-4P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of intermediates for the synthesis of discodermolides and their polyhydroxy dienyl lactone derivs. for pharmaceutical use)

RN 252342-54-4 CAPLUS

CN Phosphonium, [(2R, 3R, 4S, 5Z, 8S, 9R, 10R, 11S, 12S, 13Z)-3, 9-bis[[(1, 1dimethylethyl)dimethylsilyl]oxy]-11-[(4-methoxyphenyl)methoxy]-2,4,6,8,10,12-hexamethyl-5,13,15-hexadecatrienyl]triphenyl-, iodide (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+). Double bond geometry as shown.

I-

ANSWER 18 OF 18 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1999:694867 CAPLUS

DOCUMENT NUMBER: 132:35548

TITLE: Gram-Scale Synthesis of (+)-Discodermolide

AUTHOR(S): Smith, Amos B., III; Kaufman, Michael D.; Beauchamp,

Thomas J.; LaMarche, Matthew J.; Arimoto, Hirokazu CORPORATE SOURCE: Department of Chemistry Monell Chemical Senses Center

and Laboratory for Research on the Structure of Matter, University of Pennsylvania, PA, 19104, USA

SOURCE:

Organic Letters (1999), 1(11), 1823-1826

CODEN: ORLEF7; ISSN: 1523-7060

PUBLISHER: American Chemical Society

DOCUMENT TYPE: Journal LANGUAGE: English

A triply convergent, highly efficient second-generation synthesis of the potent antimitotic agent (+)-discodermolide has been achieved on a 1-g scale.

IT 252342-54-4P

> RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(gram-scale synthesis of (+)-discodermolide)

RN 252342-54-4 CAPLUS CN Phosphonium, [(2R,3R,4S,5Z,8S,9R,10R,11S,12S,13Z)-3,9-bis[[(1,1-dimethylethyl)dimethylsilyl]oxy]-11-[(4-methoxyphenyl)methoxy]-2,4,6,8,10,12-hexamethyl-5,13,15-hexadecatrienyl]triphenyl-, iodide (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+). Double bond geometry as shown.

• I-

REFERENCE COUNT:

THERE ARE 24 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT.

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(FILE 'HOME' ENTERED AT 08:41:40 ON 01 OCT 2007)

FILE 'REGISTRY' ENTERED AT 08:41:47 ON 01 OCT 2007

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L3 14 S L1 FULL

FILE 'CAPLUS' ENTERED AT 08:42:18 ON 01 OCT 2007

L4 18 S L3 FULL

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